

# Package ‘glyrepr’

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**Title** Representation for Glycan Compositions and Structures

**Version** 0.12.0

**Description** Computational representations of glycan compositions and structures, including details such as linkages, anomers, and substituents. Supports varying levels of monosaccharide specificity (e.g., ``Hex" or ``Gal") and ambiguous linkages. Provides robust parsing and generation of IUPAC-condensed structure strings. Optimized for vectorized operations on glycan structures, with efficient handling of duplications. As the cornerstone of the glycoverse ecosystem, this package delivers the foundational data structures that power glycomics and glycoproteomics analysis workflows.

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## Contents

as_glycan_composition . . . . .	2
as_glycan_structure . . . . .	4
available_monosaccharides . . . . .	5
available_substituents . . . . .	5
convert_to_generic . . . . .	6
count_mono . . . . .	7
fill_anomer_pos . . . . .	8
get_anomer . . . . .	9
get_anomer_pos . . . . .	9
get_mono_type . . . . .	10
get_structure_graphs . . . . .	11
get_structure_level . . . . .	12
glycan_composition . . . . .	13
glycan_structure . . . . .	14
has_linkages . . . . .	17
is_known_monosaccharide . . . . .	18
n_glycan_core . . . . .	18
possible_linkages . . . . .	20
reduce_structure_level . . . . .	21
remove_linkages . . . . .	22
remove_substituents . . . . .	22
simap . . . . .	23
smap . . . . .	25
smap2 . . . . .	26
smap_predicates . . . . .	28
smap_unique . . . . .	30
smap . . . . .	31
structure_to_iupac . . . . .	33
valid_linkages . . . . .	34
<b>Index</b>	<b>36</b>

---

as\_glycan\_composition *Convert to Glycan Composition*

---

### Description

Converts an object to a glycan composition using vctrs casting framework. This function provides a convenient way to convert various input types to `glycan_composition()`.

### Usage

```
as_glycan_composition(x)
```

**Arguments**

- x                    An object to convert to a glycan composition. Supported inputs include:
- Named integer vectors or lists of named integer vectors
  - Character vectors with composition strings (e.g., "Hex(5)HexNAc(2)")
  - glyrepr\_structure objects (counts both monosaccharides and substituents)
  - Existing glyrepr\_composition objects (returned as-is)

**Details**

This function uses the vctrs casting framework for type conversion. When converting from glycan structures, both monosaccharides and substituents are counted. Substituents are extracted from the sub attribute of each vertex in the structure. For example, a vertex with sub = "3Me" contributes one "Me" substituent to the composition.

Simple composition strings use one-letter residue codes: "H" for "Hex", "N" for "HexNAc", "F" for "dHex", "S"/"A" for "NeuAc", and "G" for "NeuGc". "E" and "L" are also accepted as linkage-specific Neu5Ac codes; they are converted to "NeuAc" with a warning because composition objects do not preserve linkage information.

**Value**

A glyrepr\_composition object.

**Examples**

```
# From a single named vector
as_glycan_composition(c(Hex = 5, HexNAc = 2))

# From a list of named vectors
as_glycan_composition(list(c(Hex = 5, HexNAc = 2), c(Hex = 3, HexNAc = 1)))

# From a character vector of Byonic composition strings
as_glycan_composition(c("Hex(5)HexNAc(2)", "Hex(3)HexNAc(1)"))

# From a character vector of simple composition strings
as_glycan_composition(c("H5N2", "H5N4S1F1"))

# From an existing composition (returns as-is)
comp <- glycan_composition(c(Hex = 5, HexNAc = 2))
as_glycan_composition(comp)

# From a glycan structure vector
strucs <- c(n_glycan_core(), o_glycan_core_1())
as_glycan_composition(strucs)
```

---

as\_glycan\_structure     *Convert to Glycan Structure Vector*

---

### Description

Convert an object to a glycan structure vector.

### Usage

```
as_glycan_structure(x)
```

### Arguments

x                    An object to convert to a glycan structure vector. Can be an igraph object, a list of igraph objects, a character vector of IUPAC-condensed strings, or an existing glyrepr\_structure object.

### Value

A glyrepr\_structure object.

### Examples

```
library(igraph)

# Convert a single igraph
graph <- make_graph(~ 1--2)
V(graph)$mono <- c("GlcNAc", "GlcNAc")
V(graph)$sub <- ""
E(graph)$linkage <- "b1-4"
graph$anomer <- "a1"
as_glycan_structure(graph)

# Convert a list of igraphs
o_glycan_vec <- o_glycan_core_1()
o_glycan_graph <- get_structure_graphs(o_glycan_vec)
as_glycan_structure(list(graph, o_glycan_graph))

# Convert a character vector of IUPAC-condensed strings
as_glycan_structure(c("GlcNAc(b1-4)GlcNAc(b1-", "Man(a1-2)GlcNAc(b1-"))
```

---

`available_monosaccharides`*Get Available Monosaccharides*

---

**Description**

This function returns a character vector of monosaccharide names of the given type. See [get\\_mono\\_type\(\)](#) for monosaccharide types.

**Usage**

```
available_monosaccharides(mono_type = "all")
```

**Arguments**

`mono_type` A character string specifying the type of monosaccharides. Can be "all", "generic", or "concrete". Default is "all".

**Value**

A character vector of monosaccharide names.

**Examples**

```
available_monosaccharides()
```

---

`available_substituents`*Available Substituents*

---

**Description**

Get the available substituents for monosaccharides.

**Usage**

```
available_substituents()
```

**Value**

A character vector.

**Examples**

```
available_substituents()
```

---

convert\_to\_generic      *Convert Monosaccharides to Generic Type*

---

## Description

This function converts monosaccharide types of monosaccharide characters, glycan compositions, or glycan structures from concrete to generic type. This is a simplified version that only supports conversion from "concrete" to "generic" monosaccharides.

## Usage

```
convert_to_generic(x)

## S3 method for class 'character'
convert_to_generic(x)

## S3 method for class 'glyrepr_structure'
convert_to_generic(x)

## S3 method for class 'glyrepr_composition'
convert_to_generic(x)
```

## Arguments

x                      Either of these objects:

- A character of monosaccharide;
- A glycan composition vector ("glyrepr\_composition" object);
- A glycan structure vector ("glyrepr\_structure" object).

## Value

A new object of the same class as x with monosaccharides converted to generic type.

## Two types of monosaccharides

There are two types of monosaccharides:

- concrete: e.g. "Gal", "GlcNAc", "Glc", "Fuc", etc.
- generic: e.g. "Hex", "HexNAc", "HexA", "HexN", etc.

For the full list of monosaccharides, use [available\\_monosaccharides\(\)](#).

**Examples**

```
# Convert character vectors
convert_to_generic(c("Gal", "GlcNAc"))

# Convert glycan compositions
comps <- glycan_composition(
  c(Gal = 5, GlcNAc = 2),
  c(Glc = 5, GalNAc = 4, Fuc = 1)
)
convert_to_generic(comps)

# Convert glycan structures
strucs <- c(n_glycan_core(), o_glycan_core_1())
convert_to_generic(strucs)
```

---

count\_mono

*Get the Number of Monosaccharides*


---

**Description**

When mono is:

- NULL (default), returns the total number of monosaccharides and substituents.
- A string, returns the number of the specified monosaccharide or substituent.

**Usage**

```
count_mono(x, mono = NULL, include_subs = FALSE)
```

```
## S3 method for class 'glyrepr_composition'
count_mono(x, mono = NULL, include_subs = FALSE)
```

```
## S3 method for class 'glyrepr_structure'
count_mono(x, mono = NULL, include_subs = FALSE)
```

**Arguments**

x	A glycan composition (glyrepr_composition) or a glycan structure (glyrepr_structure) vector.
mono	The monosaccharide or substituent to count. A character scalar. If NULL (default), return the total number of monosaccharides.
include_subs	Whether to include substituents when mono is NULL. Default is FALSE.

**Details**

When mono is "generic" (e.g. "Hex", "HexNAc"), it counts all "concrete" monosaccharides that match. For example, "Hex" will count all Glc, Man, Gal, etc. When mono is "concrete" (e.g. "Gal", "GalNAc"), NA is returned when the composition is "generic".

**Value**

A numeric vector of the same length as x.

**Examples**

```
comp <- glycan_composition(c(Gal = 1, Man = 1, GalNAc = 1))
count_mono(comp, "Hex")
count_mono(comp, "Gal")

struct <- as_glycan_structure("Gal(b1-3)GlcNAc(b1-4)Glc(a1-")
count_mono(struct, "Gal")

# Total number of monosaccharides
count_mono(comp)
```

---

fill_anomer_pos	<i>Fill Anomer Positions</i>
-----------------	------------------------------

---

**Description**

Add anomer positions to glycan structures with missing anomer position information. For example, "Gal(??-?)GalNAc(??-)" is converted to "Gal(?1-?)GalNAc(?1-)".

**Usage**

```
fill_anomer_pos(strucs)
```

**Arguments**

strucs            A [glycan\\_structure\(\)](#) vector with concrete monosaccharides.

**Details**

For anomer positions that are already specified in the input structures, this function does not modify them.

**Value**

A [glycan\\_structure\(\)](#) vector with anomer positions added where missing.

**Examples**

```
glycans <- as_glycan_structure(c(
  "Gal(??-?)GalNAc(??-)",
  "Neu5Ac(??-?)Gal(??-?)GalNAc(??-)"
))
fill_anomer_pos(glycans)
```

---

get_anomer	<i>Get the Anomeric information</i>
------------	-------------------------------------

---

**Description**

Get the Anomeric information

**Usage**

```
get_anomer(x)
```

**Arguments**

x                    A glycan structure vector (glyrepr\_structure).

**Value**

a character vector of the anomeric information.

**Examples**

```
x <- n_glycan_core()
get_anomer(x)
```

---

get_anomer_pos	<i>Get Anomer Positions</i>
----------------	-----------------------------

---

**Description**

This function returns the anomer position for concrete monosaccharide names.

**Usage**

```
get_anomer_pos(mono)
```

**Arguments**

mono                A character vector of concrete monosaccharide names.

**Value**

An integer vector of anomer positions.

**Examples**

```
get_anomer_pos(c("Gal", "Neu5Ac"))
```

---

get_mono_type	<i>Get Monosaccharide Types</i>
---------------	---------------------------------

---

### Description

This function determines the type of monosaccharides in character vectors, glycan compositions, or glycan structures. Supported types: "concrete" and "generic" (see details below).

### Usage

```
get_mono_type(x)

## S3 method for class 'character'
get_mono_type(x)

## S3 method for class 'glyrepr_structure'
get_mono_type(x)

## S3 method for class 'glyrepr_composition'
get_mono_type(x)
```

### Arguments

x Either of these objects:

- A character vector of monosaccharide names;
- A glycan composition vector ("glyrepr\_composition" object);
- A glycan structure vector ("glyrepr\_structure" object).

### Value

- For character input, returns a character vector of the same length as x.
- For glyrepr\_structure and glyrepr\_composition input, returns a character scalar.

### Two types of monosaccharides

There are two types of monosaccharides:

- concrete: e.g. "Gal", "GlcNAc", "Glc", "Fuc", etc.
- generic: e.g. "Hex", "HexNAc", "HexA", "HexN", etc.

For the full list of monosaccharides, use [available\\_monosaccharides\(\)](#).

### Special monosaccharides

Some monosaccharides are special in that they have no generic names in database or literature. For example, "Mur" is a rare monosaccharide that has no popular generic name. In glyrepr, we assign a "g" prefix to these monosaccharides as their generic names. This includes "gNeu", "gKdn", "gPse", "gLeg", "gAci", "g4eLeg", "gBac", "gKdo", "gMur". These names might only be meaningful inside glycoverse. Take care when you export results from glycoverse functions to other analysis tools.

**See Also**[convert\\_to\\_generic\(\)](#)**Examples**

```
# Character vector
get_mono_type(c("Gal", "Hex"))

# Glycan structures
get_mono_type(n_glycan_core(mono_type = "concrete"))
get_mono_type(n_glycan_core(mono_type = "generic"))

# Glycan compositions
comp <- glycan_composition(c(Glc = 2, GalNAc = 1))
get_mono_type(comp)
```

---

get\_structure\_graphs *Access Individual Glycan Structures*

---

**Description**

Extract individual glycan structure graphs from a glycan structure vector.

**Usage**

```
get_structure_graphs(x, return_list = NULL)
```

**Arguments**

x	A glycan structure vector.
return_list	If TRUE, always returns a list. If FALSE and x has a length of 1, return the igraph object directly. If not provided (default), FALSE when x has a length of 1 and TRUE otherwise.

**Value**

A list of igraph objects or an igraph object directly (see return\_list parameter).

**Examples**

```
structures <- c(o_glycan_core_1(), n_glycan_core())
get_structure_graphs(structures)
get_structure_graphs(structures)
```

---

get\_structure\_level     *Get the Structure Resolution Levels*

---

## Description

Glycan structures can have four possible levels of resolution:

- "intact": All monosaccharides are concrete (e.g. "Man", "GlcNAc"), and no linkage or anomer contains "?".
- "partial": All monosaccharides are concrete (e.g. "Man", "GlcNAc"), at least one linkage or anomer contains "?", and at least one linkage or anomer has a non-"?" annotation.
- "topological": All monosaccharides are concrete (e.g. "Man", "GlcNAc"), and all linkages and anomers are completely unknown ("??-?"/"??").
- "basic": All monosaccharides are generic (e.g. "Hex", "HexNAc").

Note that in theory you can have a glycan with generic monosaccharides with all linkages determined. For example, "Hex(b1-3)HexNAc(a1-)" is a valid glycan structure. But in reality, this is almost impossible, because linkage information is far more difficult to acquire than monosaccharide information. This kind of glycan structure is also assigned to "basic" level.

## Usage

```
get_structure_level(x)
```

## Arguments

x                    A [glycan\\_structure\(\)](#) vector.

## Value

A character scalar containing the structure level for x. If x is empty or all structures in x are NA, returns NA\_character\_.

## See Also

[has\\_linkages\(\)](#), [get\\_mono\\_type\(\)](#)

## Examples

```
glycan <- as_glycan_structure("Gal(b1-3)GalNAc(a1-")
get_structure_level(glycan)
```

---

glycan\_composition      *Create a Glycan Composition*

---

## Description

Create a glycan composition from a list of named integer vectors. Compositions can contain both monosaccharides and substituents.

## Usage

```
glycan_composition(...)
```

```
is_glycan_composition(x)
```

## Arguments

...	Named integer vectors. Names are monosaccharides or substituents, values are numbers of residues. Monosaccharides and substituents can be mixed in the same composition.
x	A list of named integer vectors.

## Details

Compositions can contain:

- Monosaccharides: either generic (e.g., "Hex", "HexNAc") or concrete (e.g., "Glc", "Gal"). All monosaccharides in a composition vector must be of the same type.
- Substituents: e.g., "Me", "Ac", "S". These can be mixed with either generic or concrete monosaccharides.

Components are automatically sorted with monosaccharides first (according to their order in the monosaccharides table), followed by substituents (according to their order in available\_substituents()). Duplicate components are automatically summed.

## Value

A glyrepr\_composition object.

## See Also

available\_monosaccharides(), available\_substituents()

## Examples

```
# A vector with one composition (generic monosaccharides)
glycan_composition(c(Hex = 5, HexNAc = 2))
# A vector with multiple compositions
glycan_composition(c(Hex = 5, HexNAc = 2), c(Hex = 5, HexNAc = 4, dHex = 2))
# Residues are reordered automatically
glycan_composition(c(HexNAc = 1, Hex = 2))
# An example for generic monosaccharides
glycan_composition(c(Hex = 2, HexNAc = 1))
# An example for concrete monosaccharides
glycan_composition(c(Glc = 2, Gal = 1))
# Compositions with substituents
glycan_composition(c(Glc = 1, S = 1))
glycan_composition(c(Hex = 3, HexNAc = 2, Me = 1, Ac = 1))
# Substituents are sorted after monosaccharides
glycan_composition(c(S = 1, Gal = 1, Ac = 1, Glc = 1))
```

---

glycan\_structure

*Create a Glycan Structure Vector*

---

## Description

glycan\_structure() creates an efficient glycan structure vector for storing and processing glycan molecular structures. The function employs hash-based deduplication mechanisms, making it suitable for glycoproteomics, glycomics analysis, and glycan structure comparison studies.

## Usage

```
glycan_structure(...)  
  
is_glycan_structure(x)
```

## Arguments

...	igraph graph objects to be converted to glycan structures, or existing glycan structure vectors. Supports mixed input of multiple objects.
x	An object to check or convert.

## Value

A glyrepr\_structure class glycan structure vector object.

## Data Structure Overview

A glycan structure vector is a `vctrs` record with an additional S3 class `glyrepr_structure`. Therefore, `sloop::s3_class()` returns the class hierarchy `c("glyrepr_structure", "vctrs_rcrd")`.

Each glycan structure must satisfy the following constraints:

### Graph Structure Requirements:

- Must be a directed graph with an outward tree structure (reducing end as root)
- Must have a graph attribute `anomer` in the format "a1" or "b1"
  - Unknown parts can be represented with "?", e.g., "?1", "a?", "??"

### Node Attributes:

- `mono`: Monosaccharide names, must be known monosaccharide types
  - Generic names: Hex, HexNAc, dHex, NeuAc, etc.
  - Concrete names: Glc, Gal, Man, GlcNAc, etc.
  - Cannot mix generic and concrete names
  - NA values are not allowed
- `sub`: Substituent information
  - Single substituent format: "xY" (x = position, Y = substituent name), e.g., "2Ac", "3S"
  - Multiple substituents separated by commas and ordered by position, e.g., "3Me,4Ac", "2S,6P"
  - No substituents represented by empty string ""

### Edge Attributes:

- `linkage`: Glycosidic linkage information in format "a/bX-Y"
  - Standard format: e.g., "b1-4", "a2-3"
  - Unknown positions allowed: "a1-?", "b?-3", "??-?"
  - Partially unknown positions: "a1-3/6", "a1-3/6/9"
  - NA values are not allowed

## Node and Edge Order

The indices of vertices and linkages in a glycan correspond directly to their order in the IUPAC-condensed string, which is printed when you print a `glycan_structure()`. For example, for the glycan `Man(a1-3)[Man(a1-6)]Man(b1-4)GlcNAc(b1-4)GlcNAc(b1-)`, the vertices are "Man", "Man", "Man", "GlcNAc", "GlcNAc", and the linkages are "a1-3", "a1-6", "b1-4", "b1-4".

## NA Support

Glycan structure vectors support NA values for representing missing or unknown structures:

- Create with `glycan_structure(NA)` or `glycan_structure(NULL)`
- Combine with valid structures: `c(struct1, NA, struct2)`
- Convert from character: `as_glycan_structure(c("Glc(a1-", NA))`
- `smap` functions skip NA elements gracefully
- `is.na()` returns TRUE for NA elements

## Naming Support

Glycan structure vectors can have names, which are preserved during operations. This is particularly useful when working with the `glymotif` package.

## As characters

One side-effect of the current implementation is that you can treat a glycan structure vector as a pure character vector of IUPAC-condensed strings. In fact, `is.character()` returns `TRUE` for a glycan structure vector, and all `stringr` functions work directly on the vector.

However, we still recommend using `as.character()` to explicitly convert to character when needed, to avoid confusion and ensure that the intended behavior is clear.

## Examples

```
library(igraph)

# Example 1: Create a simple glycan structure GlcNAc(b1-4)GlcNAc
graph <- make_graph(~ 1-+2) # Create graph with two monosaccharides
V(graph)$mono <- c("GlcNAc", "GlcNAc") # Set monosaccharide types
V(graph)$sub <- "" # No substituents
E(graph)$linkage <- "b1-4" # b1-4 glycosidic linkage
graph$anomer <- "a1" # a anomeric carbon

# Create glycan structure vector
simple_struct <- glycan_structure(graph)
print(simple_struct)

# Example 2: Use predefined glycan core structures
n_core <- n_glycan_core() # N-glycan core structure
o_core1 <- o_glycan_core_1() # O-glycan Core 1 structure

# Example 3: Create complex structure with substituents
complex_graph <- make_graph(~ 1-+2-+3)
V(complex_graph)$mono <- c("GlcNAc", "Gal", "Neu5Ac")
V(complex_graph)$sub <- c("", "", "") # Add substituents as needed
E(complex_graph)$linkage <- c("b1-4", "a2-3")
complex_graph$anomer <- "b1"

complex_struct <- glycan_structure(complex_graph)
print(complex_struct)

# Example 4: Check if object is a glycan structure
is_glycan_structure(simple_struct) # TRUE
is_glycan_structure(graph) # FALSE
```

---

has_linkages	<i>Determine if a Glycan Structure has Linkages</i>
--------------	---

---

### Description

Unknown linkages in a glycan structure are represented by "??-?". Also, a linkage can be partially known (e.g. "a?-?"). This function checks if a glycan structure has linkages, in a strict or lenient way.

### Usage

```
has_linkages(glycan, strict = FALSE)
```

### Arguments

glycan	A <a href="#">glycan_structure()</a> vector.
strict	A logical value. <ul style="list-style-type: none"><li>• If FALSE (default), a glycan is considered to have linkages if any linkage is partially known (not "??-?").</li><li>• If TRUE, a glycan is considered to have linkages only if all linkages are fully determined (no "?" in the linkage).</li></ul>

### Value

A logical vector indicating if each glycan structure has linkages.

### See Also

[remove\\_linkages\(\)](#), [possible\\_linkages\(\)](#)

### Examples

```
glycan <- o_glycan_core_1(linkage = TRUE)
has_linkages(glycan)
print(glycan)

glycan <- remove_linkages(glycan)
has_linkages(glycan)
print(glycan)

glycan <- as_glycan_structure("Gal(b1-?)GalNAc(a1-")
has_linkages(glycan)
has_linkages(glycan, strict = TRUE)
```

---

 is\_known\_monosaccharide

*Check if a Monosaccharide is Known*


---

### Description

This function checks if a vector of monosaccharide names are known.

### Usage

```
is_known_monosaccharide(mono)
```

### Arguments

mono                    A character vector of monosaccharide names.

### Value

A logical vector.

### Examples

```
is_known_monosaccharide(c("Gal", "Hex"))
is_known_monosaccharide(c("X", "Hx", "Nac"))
```

---

 n\_glycan\_core

*Example Glycan Structures*


---

### Description

Create example glycan structures for testing and demonstration. Includes **N-glycan core** and **O-glycan core 1** and **core 2**.

### Usage

```
n_glycan_core(linkage = TRUE, mono_type = "concrete")
o_glycan_core_1(linkage = TRUE, mono_type = "concrete")
o_glycan_core_2(linkage = TRUE, mono_type = "concrete")
```

### Arguments

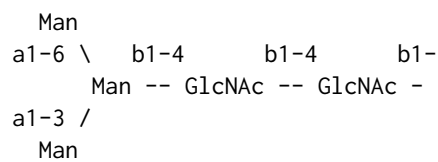
linkage                A logical indicating whether to include linkages (e.g. "b1-4"). Default is TRUE.  
 mono\_type            A character string specifying the type of monosaccharides. Can be "generic" (Hex, HexNAc, dHex, NeuAc, etc.) or "concrete" (Man, Gal, HexNAc, Fuc, etc.). Default is "concrete".

**Value**

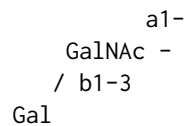
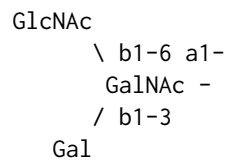
A glycan structure (igraph) object.

**N-Glycan Core**

**N-Glycans** are branched oligosaccharides that are bound, most commonly, via GlcNAc to an Asn residue of the protein backbone. A common motif of all N-glycans is the **chitobiose core**, composed of three mannose and two GlcNAc moieties, which is commonly attached to the protein backbone via GlcNAc. The mannose residue is branched and connected via  $\alpha$ 1,3- and  $\alpha$ 1,6-glycosidic linkages to the two other mannose building blocks.

**O-Glycan Core**

**O-Glycans** are highly abundant in extracellular proteins. Generally, O-glycans are extended following four major core structures: **core 1**, **core 2**, core 3, and core 4. The first two are by far the most common core structures in O-glycosylation and are found throughout the body.

**core 1:****core 2:****Examples**

```

print(n_glycan_core(), verbose = TRUE)
print(o_glycan_core_1(), verbose = TRUE)

```

---

possible\_linkages      *Generate Possible Linkages*

---

### Description

Given an obscure linkage format (having "?", e.g. "a2-?"), this function generates all possible linkages based on the format. See [valid\\_linkages\(\)](#) for details.

The ranges of possible anomers, first positions, and second positions can be specified using `anomer_range`, `pos1_range`, and `pos2_range`.

### Usage

```
possible_linkages(  
  linkage,  
  anomer_range = c("a", "b"),  
  pos1_range = 1:2,  
  pos2_range = 1:9,  
  include_unknown = FALSE  
)
```

### Arguments

`linkage`            A linkage string.  
`anomer_range`      A character vector of possible anomers. Default is `c("a", "b")`.  
`pos1_range`        A numeric vector of possible first positions. Default is `1:2`.  
`pos2_range`        A numeric vector of possible second positions. Default is `1:9`.  
`include_unknown`   A logical value. If TRUE, "?" will be included. Default is FALSE.

### Value

A character vector of possible linkages.

### See Also

[has\\_linkages\(\)](#), [remove\\_linkages\(\)](#), [valid\\_linkages\(\)](#)

### Examples

```
possible_linkages("a2-?")  
possible_linkages("??-2")  
possible_linkages("a1-3")  
possible_linkages("a?-?", pos1_range = 2, pos2_range = c(2, 3))  
possible_linkages("?1-6", include_unknown = TRUE)
```

---

`reduce_structure_level`*Reduce a Glycan Structure to a Lower Resolution Level*

---

## Description

This function reduces a glycan structure from a higher resolution level to a lower resolution level (see `get_structure_level()` for four possible levels of resolution). For example, it can reduce an "intact" structure to a "topological" structure, or a "partial" structure to a "basic" structure. One exception is that you can never reduce an "intact" structure to "partial" level, because the "partial" level is not deterministic.

## Usage

```
reduce_structure_level(x, to_level)
```

## Arguments

<code>x</code>	A <code>glycan_structure()</code> vector.
<code>to_level</code>	The resolution level to reduce to. Can be "basic" or "topological". Must be a lower resolution level than <code>x</code> ("intact" > "partial" > "topological" > "basic"). If <code>to_level</code> is the same as the structure level of <code>x</code> , the result will be the same as the input. You can use <code>get_structure_level()</code> to check the structure level of <code>x</code> .

## Details

The logic is as follows:

- If `to_level` is "topological", this function calls `remove_linkages()` to remove all linkages.
- If `to_level` is "basic", this function calls `remove_linkages()` to remove all linkages, and `convert_to_generic()` to convert all monosaccharides to generic.

## Value

A `glycan_structure()` vector reduced to the given resolution level.

## See Also

`get_structure_level()`

## Examples

```
glycan <- as_glycan_structure("Gal(b1-3)GalNAc(a1-")
reduce_structure_level(glycan, to_level = "topological")
```

---

remove_linkages	<i>Remove All Linkages from a Glycan</i>
-----------------	--

---

**Description**

This function replaces all linkages in a glycan structure with "??-", as well as the reducing end anomer with "??-".

**Usage**

```
remove_linkages(glycan)
```

**Arguments**

glycan            A glyrepr\_structure vector.

**Value**

A glyrepr\_structure vector with all linkages removed.

**Examples**

```
glycan <- o_glycan_core_1(linkage = TRUE)
glycan
remove_linkages(glycan)
```

---

remove_substituents	<i>Remove All Substituents from a Glycan</i>
---------------------	--

---

**Description**

This function replaces all substituents in a glycan structure with empty strings.

**Usage**

```
remove_substituents(glycan)
```

**Arguments**

glycan            A glyrepr\_structure vector.

**Value**

A glyrepr\_structure vector with all substituents removed.

## Examples

```
(glycan <- o_glycan_core_1())  
remove_substituents(glycan)
```

---

simap

*Map Functions Over Glycan Structure Vectors with Indices*

---

## Description

These functions apply a function to each unique structure in a glycan structure vector along with their corresponding indices, taking advantage of hash-based deduplication to avoid redundant computation. Similar to purrr imap functions, but optimized for glycan structure vectors.

## Usage

```
simap(.x, .f, ...)  
  
simap_vec(.x, .f, ..., .ptype = NULL)  
  
simap_lgl(.x, .f, ...)  
  
simap_int(.x, .f, ...)  
  
simap_dbl(.x, .f, ...)  
  
simap_chr(.x, .f, ...)  
  
simap_structure(.x, .f, ...)
```

## Arguments

<code>.x</code>	A glycan structure vector ( <code>glyrepr_structure</code> ).
<code>.f</code>	A function that takes an <code>igraph</code> object (from <code>.x</code> ) and an index/name, returning a result. Can be a function, purrr-style lambda ( <code>~ paste(.x, .y)</code> ), or a character string naming a function.
<code>...</code>	Additional arguments passed to <code>.f</code> .
<code>.ptype</code>	A prototype for the return type (for <code>simap_vec</code> ).

## Details

These functions only compute `.f` once for each unique combination of structure and corresponding index/name, then map the results back to the original vector positions. This is much more efficient than applying `.f` to each element individually when there are duplicate structures.

**IMPORTANT PERFORMANCE NOTE:** Due to the inclusion of position indices, `simap` functions have **O(total\_structures)** time complexity because each position creates a unique combination, even with identical structures.

**Alternative:** Consider `smap()` functions if position information is not required.

The index passed to `.f` is the position in the original vector (1-based). If the vector has names, the names are passed instead of indices.

### Return Types:

- `simap()`: Returns a list with the same length as `.x`
- `simap_vec()`: Returns an atomic vector with the same length as `.x`
- `simap_lgl()`: Returns a logical vector
- `simap_int()`: Returns an integer vector
- `simap_dbl()`: Returns a double vector
- `simap_chr()`: Returns a character vector
- `simap_structure()`: Returns a new glycan structure vector (`.f` must return `igraph` objects)

### Value

- `simap()`: A list
- `simap_vec()`: An atomic vector of type specified by `.ptype`
- `simap_lgl()`: Returns a logical vector
- `simap_int()`: Returns an integer vector
- `simap_dbl()`: Returns a double vector
- `simap_chr()`: Returns a character vector
- `simap_structure()`: A new `glyrepr_structure` object

### Examples

```
# Create structure vectors with duplicates
core1 <- o_glycan_core_1()
core2 <- n_glycan_core()
structures <- c(core1, core2, core1) # core1 appears twice

# Map a function that uses both structure and index
simap_chr(structures, function(g, i) paste0("Structure_", i, "_vcount_", igraph::vcount(g)))

# Use purrr-style lambda functions
simap_chr(structures, ~ paste0("Pos", .y, "_vertices", igraph::vcount(.x)))
```

**Description**

These functions apply a function to each unique structure in a glycan structure vector, taking advantage of hash-based deduplication to avoid redundant computation. Similar to purrr mapping functions, but optimized for glycan structure vectors.

**Usage**

```
smap(.x, .f, ..., .parallel = FALSE)
smap_vec(.x, .f, ..., .ptype = NULL, .parallel = FALSE)
smap_lgl(.x, .f, ..., .parallel = FALSE)
smap_int(.x, .f, ..., .parallel = FALSE)
smap_dbl(.x, .f, ..., .parallel = FALSE)
smap_chr(.x, .f, ..., .parallel = FALSE)
smap_structure(.x, .f, ..., .parallel = FALSE)
```

**Arguments**

<code>.x</code>	A glycan structure vector ( <code>glyrepr_structure</code> ).
<code>.f</code>	A function that takes an <code>igraph</code> object and returns a result. Can be a function, purrr-style lambda ( <code>~ .x\$attr</code> ), or a character string naming a function.
<code>...</code>	Additional arguments passed to <code>.f</code> .
<code>.parallel</code>	Logical; whether to use parallel processing. If <code>FALSE</code> (default), parallel processing is disabled. Set to <code>TRUE</code> to enable parallel processing.
<code>.ptype</code>	A prototype for the return type (for <code>smap_vec</code> ).

**Details**

These functions only compute `.f` once for each unique structure, then map the results back to the original vector positions. This is much more efficient than applying `.f` to each element individually when there are duplicate structures.

**Return Types:**

- `smap()`: Returns a list with the same length as `.x`
- `smap_vec()`: Returns an atomic vector with the same length as `.x`
- `smap_lgl()`: Returns a logical vector

- `smap_int()`: Returns an integer vector
- `smap_dbl()`: Returns a double vector
- `smap_chr()`: Returns a character vector
- `smap_structure()`: Returns a new glycan structure vector (`.f` must return `igraph` objects)

### Value

- `smap()`: A list
- `smap_vec()`: An atomic vector of type specified by `.ptype`
- `smap_lgl/int/dbl/chr()`: Atomic vectors of the corresponding type
- `smap_structure()`: A new `glyrepr_structure` object

### Examples

```
# Create a structure vector with duplicates
core1 <- o_glycan_core_1()
core2 <- n_glycan_core()
structures <- c(core1, core2, core1) # core1 appears twice

# Map a function that counts vertices - only computed twice, not three times
smap_int(structures, igraph::vcount)

# Map a function that returns logical
smap_lgl(structures, function(g) igraph::vcount(g) > 5)

# Use purrr-style lambda functions
smap_int(structures, ~ igraph::vcount(.x))
smap_lgl(structures, ~ igraph::vcount(.x) > 5)

# Map a function that modifies structure (must return igraph)
add_vertex_names <- function(g) {
  if (!("name" %in% igraph::vertex_attr_names(g))) {
    igraph::set_vertex_attr(g, "name", value = paste0("v", seq_len(igraph::vcount(g))))
  } else {
    g
  }
}
smap_structure(structures, add_vertex_names)
```

### Description

These functions apply a function to each unique structure combination in two glycan structure vectors, taking advantage of hash-based deduplication to avoid redundant computation. Similar to `purrr` `map2` functions, but optimized for glycan structure vectors.

**Usage**

```

smap2(.x, .y, .f, ..., .parallel = FALSE)

smap2_vec(.x, .y, .f, ..., .ptype = NULL, .parallel = FALSE)

smap2_lgl(.x, .y, .f, ..., .parallel = FALSE)

smap2_int(.x, .y, .f, ..., .parallel = FALSE)

smap2_dbl(.x, .y, .f, ..., .parallel = FALSE)

smap2_chr(.x, .y, .f, ..., .parallel = FALSE)

smap2_structure(.x, .y, .f, ..., .parallel = FALSE)

```

**Arguments**

<code>.x</code>	A glycan structure vector ( <code>glyrepr_structure</code> ).
<code>.y</code>	A vector of the same length as <code>.x</code> , or length 1 (will be recycled).
<code>.f</code>	A function that takes an <code>igraph</code> object (from <code>.x</code> ) and a value (from <code>.y</code> ) and returns a result. Can be a function, purrr-style lambda ( <code>~ .x + .y</code> ), or a character string naming a function.
<code>...</code>	Additional arguments passed to <code>.f</code> .
<code>.parallel</code>	Logical; whether to use parallel processing. If <code>FALSE</code> (default), parallel processing is disabled. Set to <code>TRUE</code> to enable parallel processing. See examples in <a href="#">smap</a> for how to set up and use parallel processing.
<code>.ptype</code>	A prototype for the return type (for <code>smap2_vec</code> ).

**Details**

These functions only compute `.f` once for each unique combination of structure and corresponding `.y` value, then map the results back to the original vector positions. This is much more efficient than applying `.f` to each element pair individually when there are duplicate structure-value combinations.

**NA Handling:** NA elements in `.x` are preserved in the output - the function is not applied to NA positions, and the corresponding results are set to NA.

**Return Types:**

- `smap2()`: Returns a list with the same length as `.x`
- `smap2_vec()`: Returns an atomic vector with the same length as `.x`
- `smap2_lgl()`: Returns a logical vector
- `smap2_int()`: Returns an integer vector
- `smap2_dbl()`: Returns a double vector
- `smap2_chr()`: Returns a character vector
- `smap2_structure()`: Returns a new glycan structure vector (`.f` must return `igraph` objects)

**Value**

- `smap2()`: A list
- `smap2_vec()`: An atomic vector of type specified by `.ptype`
- `smap2_lgl/int/dbl/chr()`: Atomic vectors of the corresponding type
- `smap2_structure()`: A new `glyrepr_structure` object

**Examples**

```
# Create structure vectors with duplicates
core1 <- o_glycan_core_1()
core2 <- n_glycan_core()
structures <- c(core1, core2, core1) # core1 appears twice
weights <- c(1.0, 2.0, 1.0) # corresponding weights

# Map a function that uses both structure and weight
smap2_dbl(structures, weights, function(g, w) igraph::vcount(g) * w)

# Use purrr-style lambda functions
smap2_dbl(structures, weights, ~ igraph::vcount(.x) * .y)

# Test with recycling (single weight for all structures)
smap2_dbl(structures, 2.5, ~ igraph::vcount(.x) * .y)

# Map a function that modifies structure based on second argument
# This example adds a graph attribute instead of modifying topology
add_weight_attr <- function(g, weight) {
  igraph::set_graph_attr(g, "weight", weight)
}
weights_to_add <- c(1.5, 2.5, 1.5)
smap2_structure(structures, weights_to_add, add_weight_attr)
```

---

`smap_predicates`*Test Predicates on Glycan Structure Vectors*

---

**Description**

These functions test predicates on unique structures in a glycan structure vector, taking advantage of hash-based deduplication to avoid redundant computation. Similar to `purrr` predicate functions, but optimized for glycan structure vectors.

**Usage**`ssome(.x, .p, ...)``severy(.x, .p, ...)``snone(.x, .p, ...)`

## Arguments

<code>.x</code>	A glycan structure vector ( <code>glyrepr_structure</code> ).
<code>.p</code>	A predicate function that takes an <code>igraph</code> object and returns a logical value. Can be a function, purrr-style lambda ( <code>~ .x\$attr</code> ), or a character string naming a function.
<code>...</code>	Additional arguments passed to <code>.p</code> .

## Details

These functions only evaluate `.p` once for each unique structure, making them much more efficient than applying `.p` to each element individually when there are duplicate structures.

### Return Values:

- `ssome()`: Returns TRUE if at least one unique structure satisfies the predicate
- `severy()`: Returns TRUE if all unique structures satisfy the predicate
- `snone()`: Returns TRUE if no unique structures satisfy the predicate

## Value

A single logical value.

## Examples

```
# Create a structure vector with duplicates
core1 <- o_glycan_core_1()
core2 <- n_glycan_core()
structures <- c(core1, core2, core1) # core1 appears twice

# Test if some structures have more than 5 vertices
ssome(structures, function(g) igraph::vcount(g) > 5)

# Test if all structures have at least 3 vertices
severy(structures, function(g) igraph::vcount(g) >= 3)

# Test if no structures have more than 20 vertices
snone(structures, function(g) igraph::vcount(g) > 20)

# Use purrr-style lambda functions
ssome(structures, ~ igraph::vcount(.x) > 5)
severy(structures, ~ igraph::vcount(.x) >= 3)
snone(structures, ~ igraph::vcount(.x) > 20)
```

---

`smap_unique`*Apply Function to Unique Structures Only*

---

### Description

Apply a function only to the unique structures in a glycan structure vector, returning results in the same order as the unique structures appear. This is useful when you need to perform expensive computations but only care about unique results.

### Usage

```
smap_unique(.x, .f, ..., .parallel = FALSE)
```

### Arguments

<code>.x</code>	A glycan structure vector ( <code>glyrepr_structure</code> ).
<code>.f</code>	A function that takes an <code>igraph</code> object and returns a result. Can be a function, purrr-style lambda ( <code>~ .x\$attr</code> ), or a character string naming a function.
<code>...</code>	Additional arguments passed to <code>.f</code> .
<code>.parallel</code>	Logical; whether to use parallel processing. If <code>FALSE</code> (default), parallel processing is disabled. Set to <code>TRUE</code> to enable parallel processing. See examples in <a href="#">smap</a> for how to set up and use parallel processing.

### Value

A list with results for each unique structure, named by their hash codes.

### Examples

```
# Create a structure vector with duplicates
core1 <- o_glycan_core_1()
structures <- c(core1, core1, core1) # same structure 3 times

# Only compute once for the unique structure
unique_results <- smap_unique(structures, igraph::vcount)
length(unique_results) # 1, not 3

# Use purrr-style lambda
unique_results2 <- smap_unique(structures, ~ igraph::vcount(.x))
length(unique_results2) # 1, not 3
```

---

smap	<i>Map Functions Over Glycan Structure Vectors and Multiple Arguments</i>
------	---

---

## Description

These functions apply a function to each unique structure in a glycan structure vector along with corresponding elements from multiple other vectors, taking advantage of hash-based deduplication to avoid redundant computation. Similar to purrr pmap functions, but optimized for glycan structure vectors.

## Usage

```
smap(.l, .f, ..., .parallel = FALSE)

smap_vec(.l, .f, ..., .ptype = NULL, .parallel = FALSE)

smap_lgl(.l, .f, ..., .parallel = FALSE)

smap_int(.l, .f, ..., .parallel = FALSE)

smap_dbl(.l, .f, ..., .parallel = FALSE)

smap_chr(.l, .f, ..., .parallel = FALSE)

smap_structure(.l, .f, ..., .parallel = FALSE)
```

## Arguments

.l	A list where the first element is a glycan structure vector ( <code>glyrepr_structure</code> ) and the remaining elements are vectors of the same length or length 1 (will be recycled).
.f	A function that takes an <code>igraph</code> object (from first element of <code>.l</code> ) and values from other elements, returning a result. Can be a function, purrr-style lambda ( <code>~ .x + .y + .z</code> ), or a character string naming a function.
...	Additional arguments passed to <code>.f</code> .
.parallel	Logical; whether to use parallel processing. If <code>FALSE</code> (default), parallel processing is disabled. Set to <code>TRUE</code> to enable parallel processing. See examples in <a href="#">smap</a> for how to set up and use parallel processing.
.ptype	A prototype for the return type (for <code>smap_vec</code> ).

## Details

These functions only compute `.f` once for each unique combination of structure and corresponding values from other vectors, then map the results back to the original vector positions.

**NA Handling:** NA elements in the first argument (glycan structure vector) are preserved in the output.

**Time Complexity Performance:**

Performance scales with unique combinations of all arguments rather than total vector length. When argument vectors are highly redundant, performance approaches  $O(\text{unique\_structures})$ . Scaling factor shows time increase when vector size increases 20x.

**Return Types:**

- `smap()`: Returns a list with the same length as the input vectors
- `smap_vec()`: Returns an atomic vector with the same length as the input vectors
- `smap_lgl()`: Returns a logical vector
- `smap_int()`: Returns an integer vector
- `smap_dbl()`: Returns a double vector
- `smap_chr()`: Returns a character vector
- `smap_structure()`: Returns a new glycan structure vector (.f must return igraph objects)

**Value**

- `smap()`: A list
- `smap_vec()`: An atomic vector of type specified by .ptype
- `smap_lgl/int/dbl/chr()`: Atomic vectors of the corresponding type
- `smap_structure()`: A new glyrepr\_structure object

**Examples**

```
# Create structure vectors with duplicates
core1 <- o_glycan_core_1()
core2 <- n_glycan_core()
structures <- c(core1, core2, core1) # core1 appears twice
weights <- c(1.0, 2.0, 1.0) # corresponding weights
factors <- c(2, 3, 2) # corresponding factors

# Map a function that uses structure, weight, and factor
smap_dbl(list(structures, weights, factors),
         function(g, w, f) igraph::vcount(g) * w * f)

# Use purrr-style lambda functions
smap_dbl(list(structures, weights, factors), ~ igraph::vcount(..1) * ..2 * ..3)

# Test with recycling
smap_dbl(list(structures, 2.0, 3), ~ igraph::vcount(..1) * ..2 * ..3)
```

---

structure_to_iupac	<i>Convert Glycan Structure to IUPAC-like Sequence</i>
--------------------	--

---

### Description

Convert a glycan structure to a sequence representation in the form of mono(linkage)mono, with branches represented by square brackets []. The backbone is chosen as the longest path, and for branches, linkages are ordered lexicographically with smaller linkages on the backbone.

### Usage

```
structure_to_iupac(glycan)
```

### Arguments

glycan            A glyrepr\_structure vector.

### Value

A character vector representing the IUPAC sequences.

### Sequence Format

The sequence follows the format mono(linkage)mono, where:

- mono: monosaccharide name with optional substituents (e.g., Glc, GlcNAc, Glc3Me)
- linkage: glycosidic linkage (e.g., b1-4, a1-3)
- Branches are enclosed in square brackets []
- Substituents are appended directly to monosaccharide names (e.g., Glc3Me for Glc with 3Me substituent)

### Backbone Selection

The backbone is selected as the longest path in the tree. For branches, the same rule applies recursively.

### Linkage Comparison

Linkages are compared lexicographically:

1. First by anomeric configuration: ? > b > a
2. Then by first position: ? > numbers (numerically)
3. Finally by second position: ? > numbers (numerically)

Smaller linkages are placed on the backbone, larger ones in branches.

**Examples**

```

# Simple linear structure
structure_to_iupac(o_glycan_core_1())

# Branched structure
structure_to_iupac(n_glycan_core())

# Structure with substituents
graph <- igraph::make_graph(~ 1--+2)
igraph::V(graph)$mono <- c("Glc", "GlcNAc")
igraph::V(graph)$sub <- c("3Me", "6Ac")
igraph::E(graph)$linkage <- "b1-4"
graph$anomer <- "a1"
glycan <- glycan_structure(graph)
structure_to_iupac(glycan) # Returns "GlcNAc6Ac(b1-4)Glc3Me(a1-"

# Vectorized structures
structs <- c(o_glycan_core_1(), n_glycan_core())
structure_to_iupac(structs)

```

---

valid\_linkages

*Check if Linkages are Valid*


---

**Description**

Valid linkages are in the form of "a1-2", "b1-4", "a?-1", etc. Specifically, the pattern is xy-z:

- x: the anomer, either "a", "b", or "?".
- y: the first position, either "1", "2" or "?".
- z: the second position, either a 1-9 digit or "?". Can also be multiple positions separated by "/", e.g. "1/2/3". "?" could not be used with "/".

**Usage**

```
valid_linkages(linkages)
```

**Arguments**

linkages            A character vector of linkages.

**Value**

A logical vector.

**Examples**

```
# Valid linkages
valid_linkages(c("a1-2", "?1-4", "a?-1", "b?-?", "??-?", "a1/2-3"))

# Invalid linkages
valid_linkages(c("a1-2/?", "1-4", "a/b1-2", "c1-2", "a9-1"))
```

# Index

as\_glycan\_composition, 2  
as\_glycan\_structure, 4  
available\_monosaccharides, 5  
available\_monosaccharides(), 6, 10  
available\_substituents, 5  
  
convert\_to\_generic, 6  
convert\_to\_generic(), 11, 21  
count\_mono, 7  
  
fill\_anomer\_pos, 8  
  
get\_anomer, 9  
get\_anomer\_pos, 9  
get\_mono\_type, 10  
get\_mono\_type(), 5, 12  
get\_structure\_graphs, 11  
get\_structure\_level, 12  
get\_structure\_level(), 21  
glycan\_composition, 13  
glycan\_composition(), 2  
glycan\_structure, 14  
glycan\_structure(), 8, 12, 15, 17, 21  
  
has\_linkages, 17  
has\_linkages(), 12, 20  
  
is\_glycan\_composition  
    (glycan\_composition), 13  
is\_glycan\_structure (glycan\_structure),  
    14  
is\_known\_monosaccharide, 18  
  
n\_glycan\_core, 18  
  
o\_glycan\_core\_1 (n\_glycan\_core), 18  
o\_glycan\_core\_2 (n\_glycan\_core), 18  
  
possible\_linkages, 20  
possible\_linkages(), 17  
  
reduce\_structure\_level, 21  
  
remove\_linkages, 22  
remove\_linkages(), 17, 20, 21  
remove\_substituents, 22  
  
severy (smap\_predicates), 28  
simap, 23  
simap\_chr (simap), 23  
simap\_dbl (simap), 23  
simap\_int (simap), 23  
simap\_lgl (simap), 23  
simap\_structure (simap), 23  
simap\_vec (simap), 23  
smap, 25, 27, 30, 31  
smap2, 26  
smap2\_chr (smap2), 26  
smap2\_dbl (smap2), 26  
smap2\_int (smap2), 26  
smap2\_lgl (smap2), 26  
smap2\_structure (smap2), 26  
smap2\_vec (smap2), 26  
smap\_chr (smap), 25  
smap\_dbl (smap), 25  
smap\_int (smap), 25  
smap\_lgl (smap), 25  
smap\_predicates, 28  
smap\_structure (smap), 25  
smap\_unique, 30  
smap\_vec (smap), 25  
snone (smap\_predicates), 28  
spmap, 31  
spmap\_chr (spmap), 31  
spmap\_dbl (spmap), 31  
spmap\_int (spmap), 31  
spmap\_lgl (spmap), 31  
spmap\_structure (spmap), 31  
spmap\_vec (spmap), 31  
ssome (smap\_predicates), 28  
structure\_to\_iupac, 33  
  
valid\_linkages, 34

`valid_linkages()`, [20](#)